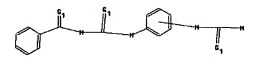
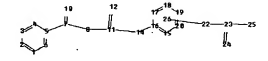
10/616,959

FILE 'HOME' ENTERED AT 10:21:01 ON 11 OCT 2006

=> file reg

=>Uploading C:\Program Files\Stnexp\Queries\10616959.str





chain nodes :

7 8 10 11 12 14 22 23 24

ring nodes :

1 2 3 4 5 6 15 16 17 18 19 20

ring/chain nodes :

25

chain bonds :

5-7 7-8 7-10 8-11 11-12 11-14 14-16 22-23 23-24 23-25

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 15-16 15-20 16-17 17-18 18-19 19-20

exact/norm bonds :

7-8 7-10 8-11 11-12 11-14 14-16 22-23 23-24 23-25

exact bonds :

5-7

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 15-16 15-20 16-17 17-18 18-19 19-20

isolated ring systems :

containing 1 : 15 :

G1:0,S

Match level:

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 10:CLASS

11:CLASS 12:CLASS 14:CLASS 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom 20:Atom

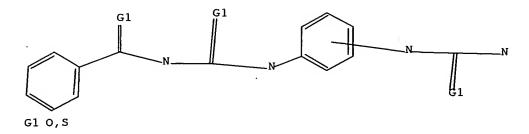
22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:Atom

L1 STRUCTURE UPLOADED

=> dis 11

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> file caplus

=> s 13

L4 39 L3

=> dis 15 1-34 bib abs hitstr

- L5 ANSWER 1 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 2002:506002 CAPLUS Full-text
- DN 137:370017
- TI A facile synthesis of p-Bis(4-thiazolidinon-3-yl)phenylenes and related systems
- AU Abdel-Megid, M.; Awas, M. A. A.
- CS Chemistry Department, Faculty of Education, Ain-Shams University, Cairo, Egypt
- SO Heterocyclic Communications (2002), 8(2), 161-168 CODEN: HCOMEX; ISSN: 0793-0283
- PB Freund Publishing House Ltd.
- DT Journal
- LA English
- OS CASREACT 137:370017

GI

$$C1$$
 $NO2$
 $NO2$
 $NO2$
 $NO2$
 $NO2$
 $NO2$
 $NO2$
 $NO2$

AB P-Bis(4-thiazolidinon-3-yl)phenylenes, e.g., I and II, were synthesized by cycloaddn. of thioglycolic acid with Schiff bases of p-phenylenediamine or by treatment of p-bis(thioureido)phenylenes with Et chloroacetate. Reactions of hydrazines, hydroxylamine, acetamidine and N-phenylthiourea with I and II were reported. Some of the new compds. were tested for their effect on cellobiase, produced by thermophilic fungi.

IT 493026-96-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of p-bis(4-thiazolidinon-3-yl)phenylenes and related systems and their effect on fungal cellobiase)

RN 493026-96-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[4-chloro- (9CI) (CA INDEX NAME)

RE.CNT 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 2 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2002:437635 CAPLUS Full-text

DN 138:137007

TI Phase transfer catalytic synthesis of phenylene-1,4-bis-aroyl(aryloxyacetyl)thiourea derivatives

AU Deng, Hong-tao; Ye, Wen-fa; Wang, Yan-gang

CS Department of Chemistry, Central China Normal University, Wuhan, 430079, Peop. Rep. China

SO Huazhong Shifan Daxue Xuebao Zirankexueban (2002), 36(1), 58-60 CODEN: HDZKEL; ISSN: 1000-1190

PB Huazhong Shifan Daxue Xuebao Bianjibu

DT Journal

LA Chinese

OS CASREACT 138:137007

AB Using p-phenylenediamine and arom. acid or aryloxyacetic acid as raw materials, PEG-600 as catalyst, ten new phenylene-1,4-bis-aroyl(aryloxyacetyl)thiourea derivs. have been synthesized by solid-liquid phase transfer catalysis. Title compds. showed plant growth regulator activities.

RN 331862-02-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3-nitro-(9CI) (CA INDEX NAME)

RN 493026-92-9 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3-methyl- (9CI) (CA INDEX NAME)

RN 493026-94-1 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[4-methoxy- (9CI) (CA INDEX NAME)

RN 493026-96-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[4-chloro-(9CI) (CA INDEX NAME)

RN 493026-98-5 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[4-nitro- (9CI) (CA INDEX NAME)

RN 493027-01-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3,5-dinitro-(9CI) (CA INDEX NAME)

L5 ANSWER 3 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2001:76696 CAPLUS Full-text

DN 134:266079

TI Phase transfer catalyzed synthesis of arene-bis-aroyl thiourea derivatives

AU Zhang, You-Ming; Wei, Tai-Bao; Gao, Li-Ming

CS Department of Chemistry, Northwest Normal University, Lanzhou, 730 070, Peop. Rep. China

SO Indian Journal of Chemistry, Section B: Organic Chemistry Including Medicinal Chemistry (2000), 39B(9), 700-702 CODEN: IJSBDB; ISSN: 0376-4699

PB National Institute of Science Communication, CSIR

DT Journal

LA English

OS CASREACT 134:266079

AB Reaction of 4.5 mmol arene diamines [1,2- and 1,4-(H2N)2C6H4, 4-H2NC6H4C6H4NH2-4, 4-H2N-3-MeC6H4C6H4Me-3-NH2-4] with 10 mmol aroyl chloride RCOCl (R = Ph, m-O2NC6H4, 2-furyl) and 15 mmol ammonium thiocyanate in 25 mL CH2Cl2 under the conditions of solid-liquid phase transfer catalysis using 3% (with respect to NH4SCN) polyethylene-glycol- 600 (PEG-600) as the catalyst furnishes 12 arene-bis-aroyl thioureas in good to excellent (86-98%) yields. E.g., reaction of BzCl with 1,4-(H2N)2C6H4 and NH4SCN in CH2Cl2 containing PEG-600 gave 98% p-BzNHC(S)NHC6H4NHC(S)NHBz. The products were characterized by anal. and spectral (IR and 1H NMR) data.

IT 70110-39-3P 87874-16-6P 331862-02-3P

331862-04-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (phase-transfer carbamoylation of in-situ formed aroyl isothiocyanates with arene diamines)

RN 70110-39-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

RN 87874-16-6 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

RN 331862-02-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3-nitro-(9CI) (CA INDEX NAME)

RN 331862-04-5 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis[3-nitro-(9CI) (CA INDEX NAME)

RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 4 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1999:463015 CAPLUS Full-text

DN 131:151789

TI Optical recording medium including carbonylthiourea derivative and its

coloring method

IN Aoki, Izuo; Sakamoto, Yasuko; Ohsawa, Michiyo

PA Nippon Soda Co., Ltd., Japan

SO Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

P.F	ATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
	-					
PI JE	2 11198547	A2	19990727	JP 1998-17831	19980114 <	
PRAI JE	1998-17831		19980114			

AB The medium contains a coloring (preferably a leuco) dye and a compd. bearing NHCSNHCO group, and is colored by pressure application, heating, or by photoirradn.

IT 87874-16-6

RL: MOA (Modifier or additive use); PEP (Physical, engineering or chemical process); TEM (Technical or engineered material use); PROC (Process); USES (Uses)

(developer; optical recording medium including fluoran-type leuco dye and carbonylthiourea derivative as developer)

RN 87874-16-6 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

- L5 ANSWER 5 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1998:104912 CAPLUS Full-text
- DN 128:154466
- TI Synthesis, characterization and electrical conductivity of polyesters, polyamides and doped polymers
- AU Bhatt, Vasishta D.; Ray, Arabinda
- CS Department of Chemistry, S.P. University, Vallabh Vidyanagar, 388120, India
- SO Synthetic Metals (1998), 92(2), 115-120 CODEN: SYMEDZ; ISSN: 0379-6779
- PB Elsevier Science S.A.
- DT Journal
- LA English
- AB Polyamides and polyesters contg. azomethyne linkages were prepd. by condensation from thioamide monomers and acid chlorides and from Schiff's bases and terephthalic acid chloride and isophthalic acid chloride, resp. The elec. conductivity of the resulting conducting polymers was studied using simple PPP [PPP] calcns. and exptl. measurements. The UV spectra of monomers and polymers indicate π π * transitions, however, no correlation could be obtained of this transition and conductivity A reasonably good correlations was obtained between the conductivity of the polymers and the frontier electron d. at the C* atom, from the LUMO [LUMO] and the next higher unoccupied orbital of the repeating unit. Upon doping with Ag, the elec. conductivity all polymers increased significantly, which is attributed to

contributions of all unoccupied orbitals of adjacent repeating units to the C* atom.

IT 70113-14-3P 202803-51-8P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)

(preparation and electronic structure and elec. conductivity of undoped and silver-doped azomethyne group-containing polyester and thio group containing

polyamide conducting polymers)

RN 70113-14-3 CAPLUS

CN Poly(iminocarbonothioylimino-1,4-phenyleneiminocarbonothioyliminocarbonyl-1,3-phenylenecarbonyl) (9CI) (CA INDEX NAME)

RN 202803-51-8 CAPLUS

CN Poly(iminocarbonothioylimino-1,4-phenyleneiminocarbonothioyliminocarbonyl-1,4-phenylenecarbonyl) (9CI) (CA INDEX NAME)

RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 6 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1995:526587 CAPLUS Full-text

DN 122:267065

TI Compounds containing two thiourea groups and their use in near-infrared absorbers and heat-blocking materials

IN Hayasaka, Hideki; Takano, Toshiyuki; Satake, Toshimi

PA Nippon Paper Industries Co., Ltd., Japan

SO Eur. Pat. Appl., 47 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

T 1 TI 4 *	ON I I				
	PATENT NO.	KIND DATE		APPLICATION NO.	DATE
ΡI	EP 611754	A 1	19940824	· EP 1994-301189	19940218 <
	EP 611754	B1	19980422		
	R: DE, FR, IT				
	JP 06299139	A2	19941025	JP 1993-199664	19930811 <
	JP 3603315	B2	20041222 .		
	AU 9455219	A1	19940825	AU 1994-55219	19940218 <
	AU 683031	B2	19971030		
	US 5723075	Α	19980303	US 1996-634126	19960419 <

PRAI JP 1993-30954 A 19930219 JP 1993-199664 A 19930811 US 1994-197948 B1 19940217

os MARPAT 122:267065

Thiourea derivs. RNHCSNHZ1AZ2NHCSNHR and RNHCSNHZ3NHCSNHR (R = alkyl, aralkyl, aryl, acyl, alkenyl, alkoxycarbonyl, etc.; A = CH2, CH2CH2, S, O, CONH, NH, etc.; Z1-2 = 1,4-phenylene optionally substituted by alkyl, nitro, cyano, and/or halo groups; Z3 = arylene or substituted arylene) having high decomposition temps. are prepared and used with Cu compds. in resin moldings which absorb near-IR radiation. Reacting PhCH2NCS with bis(4-aminophenyl)methane gave (PhCH2NHCSNH-p-C6H4)2CH2 (decomposition temperature 210.5°) which was mixed with CU stearate and polystyrene at 190° and extruded to give a near-IR absorber.

IT 162781-28-4P

RL: IMF (Industrial manufacture); POF (Polymer in formulation); PRP (Properties); PREP (Preparation); USES (Uses)

(preparation and use as heat-resistant near-IR absorbers)

RN 162781-28-4 CAPLUS

CN Benzamide, N,N'-[(2,5-dimethyl-4,1-phenylene)bis(iminocarbonothioyl)]bis-(9CI) (CA INDEX NAME)

L5 ANSWER 7 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1991:134953 CAPLUS Full-text

DN 114:134953

TI Mononuclear and binuclear nickel(II) chelates of some thiocarbamides

AU Abu El-Reash, G. M.; Taha, F. I.; Badr, G. E.

CS Fac. Sci., Mansoura Univ., Mansoura, Egypt

SO Bulletin de la Societe Chimique de France (1990), (May-June), 387-90

CODEN: BSCFAS; ISSN: 0037-8968

DT Journal

LA English

Ni(II) complexes of known thiocarbamides were prepd. from the reaction of benzoylisothiocyanate with 2-aminopyridine, 3-aminopyridine (H2L), 2,3-diaminopyridine, 2,6-diaminopyridine, o-phenylenediamine (H2L1), and ethylenediamine (H2L2). These complexes characterized by elemental analyses, molar conductivities, magnetic moments and spectral (visible, IR) measurements. IR spectra show that the ligands behave as a diamionic or neutral tetradentate ligand, as monoanionic tridentate ligand or as neutral bidentate ligand. Magnetic measurements and spectral anal. suggest that Ni(L1) and [Ni(L2)]2 are square plane, Ni(HL).2H2O is associated with a tetrahedral-square planar isomerism and the other complexes are tetragonally distorted octahedra. All the complexes are nonelectrolytes.

IT 87874-16-6

RL: PRP (Properties)

(IR spectrum of) 87874-16-6 CAPLUS

RN 87874-16-6 CAPLUS
CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CAINDEX NAME)

L5 ANSWER 8 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1990:244915 CAPLUS Full-text

DN 112:244915

TI Complexes of copper(II) with some new thiocarbamide derivatives

AU Abu El-Reash, Gaber M.; Taha, Fatma I.; Badr, Gamila

CS Fac. Sci., Mansoura Univ., Mansoura, Egypt

SO Transition Metal Chemistry (Dordrecht, Netherlands) (1990), 15(2), 116-19

CODEN: TMCHDN; ISSN: 0340-4285

DT Journal

LA English

AB A new series of thiocarbamides was prepd. by the reaction of benzoylisothiocyanate with 2-aminopyridine, 3-aminopyridine, 2,3-diaminopyridine, 2,6-diaminopyridine, o-phenylenediamine, p-phenylenediamine, and ethylenediamine. The Cu(II) complexes of these ligands were isolated and characterized by elemental analyses, molar conductivities, magnetic moments and spectral (visible, IR) measurements. IR spectra show that the ligands behave as dianionic or neutral tetradentates or as monoanionic, or neutral bidentates. [Cu(HL)Cl]2 (H2L = RNHCSNHBz (R = 2-pyridyl)) and Cu(H2L1)Cl2 (H2L1 = R1(NHCSNHBz)2 (R1 = 2,6-pyridinediyl) are diamagnetic and the other complexes have normal magnetic moment at room temperature Electronic spectral analyses show that Cu2(L1)(OAc)2 is planar and the other complexes are tetragonally distorted octahedral. All the complexes are nonelectrolytes.

IT 70110-39-3P 87874-16-6P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and IR spectrum of)

RN 70110-39-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

RN 87874-16-6 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

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L5
     ANSWER 9 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN
     1989:553377 CAPLUS Full-text
AN
DN
     111:153377
ΤI
     Benzoylurea derivatives as insecticides and acaricides and their
    preparation
     Kariya, Akinori; Nanjo, Katsumi; Katsurayama, Takayoshi
IN
    Agro-Kanesho Co., Ltd., Japan
PΑ
so
     Jpn. Kokai Tokkyo Koho, 6 pp.
    CODEN: JKXXAF
DΤ
     Patent
LА
     Japanese
FAN.CNT 1
     PATENT NO.
                         KIND
                                DATE
                                           APPLICATION NO.
                                                                  DATE
                                            -----
                                                                   -----
PΙ
    JP 01034953
                         A2
                                19890206
                                           JP 1987-190899
                                                                   19870730 <--
PRAI JP 1987-190899
                                19870730
OS
    MARPAT 111:153377
GΙ
```

$$X_{R1}$$
 CONHCONH X_{n} NHCONR² X_{m}

AB

1; R2 = lower alkyl, alkenyl; Y = H, halo, lower alkyl, alkoxy, etc.; m = 0-3), useful as insecticides and acaricides, were prepared A mixture of N-(3fluoro-4-aminophenyl)-N'-(4-chlorophenyl)-N'-propylurea and 2,6difluorobenzoyl isocyanate in ether was stirred at room temperature for 30 min to give I (R = R1 = F, Xn = H, R2 = Pr, Ym = 4-C1) (II). At 500 ppm, II gave complete control of Plutella xylostella larvae. A wettable powder containing II 40, SiO2 2, clay 53, Na alkylbenzenesulfonate 2, and naphthalenesulfonic acid formalin condensation product 3 parts was prepared IT 122815-63-8P 122815-64-9P 122815-65-0P 122815-66-1P 122815-67-2P 122815-68-3P 122815-69-4P 122815-70-7P 122815-71-8P 122815-72-9P 122815-73-0P 122815-74-1P 122815-75-2P 122815-76-3P 122815-77-4P 122815-78-5P 122815-79-6P 122815-80-9P 122815-81-0P 122815-82-1P 122815-83-2P 122815-84-3P 122815-85-4P 122815-86-5P 122815-87-6P 122815-88-7P 122815-89-8P 122829-04-3P RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as insecticide and acaricide) RN 122815-63-8 CAPLUS CN Benzamide, N-[[[4-[[(4-chlorophenyl)propylamino]carbonyl]amino]-2-

The title compds. I (R = halo; R1 = halo, H; X = H, halo, lower alkyl; n = 0,

fluorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-64-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[(4-chlorophenyl)propylamino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \circ & \circ & \\ & \downarrow & \\ &$$

RN 122815-65-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(2-chlorophenyl)propylamino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-66-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(3,4-dichlorophenyl)propylamino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

$$\bigcap_{C1} \bigcap_{NH} \bigcap_{H-NH} \bigcap_{NH-C-N} \bigcap_{NH-C-N} \bigcap_{C1} \bigcap_{C$$

RN 122815-67-2. CAPLUS

CN Benzamide, 2-chloro-N-[[[2-fluoro-4-[[[propyl[4-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-68-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)ethylamino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-69-4 CAPLUS

CN Benzamide, N-[[[4-[[(4-chlorophenyl)ethylamino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro-(9CI) (CA INDEX NAME)

RN 122815-70-7 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-chlorophenyl)amino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

RN 122815-71-8 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-72-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

$$\bigcup_{l=1}^{0}\bigcup_{l=NH-}^{0}\bigcup_{l=NH-}^{0}\bigcup_{l=NH-}^{0}\bigcup_{l=NH-}^{0}\bigcup_{l=1}^{0}\bigcup$$

RN 122815-73-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-difluoro-4-[[(phenylpropylamino)carbonyl]amin o]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-74-1 CAPLUS

CN Benzamide, N-[[[2,5-difluoro-4-[[[(4-methylphenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-75-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-difluoro-4-[[[(4-methylphenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

$$\bigcup_{C1}^{O} \bigcup_{NH-\stackrel{U}{U}-NH}^{O} \bigcup_{NH-\stackrel{U}{U}-NH}^{F} \bigcup_{F}^{Pr-n} \bigcup_{Me}^{NH-\stackrel{O}{U}-NH} \bigcup_{NH-\stackrel{U}{U}-NH}^{Pr-n} \bigcup_{Me}^{NH-\stackrel{O}{U}-NH} \bigcup_{NH-\stackrel{U}{U}-NH}^{Pr-n} \bigcup_{N$$

RN 122815-76-3 CAPLUS

CN Benzamide, N-[[[2,5-difluoro-4-[[[(4-methoxyphenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-77-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-difluoro-4-[[[(4-methoxyphenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-78-5 CAPLUS

CN Benzamide, N-[[[2,5-difluoro-4-[[[[4-(1-methylethyl)phenyl]propylamino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-79-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-difluoro-4-[[[[4-(1-methylethyl)phenyl]propylamino]carbonyl]amino]phenyl]amino]carbonyl](9CI) (CA INDEX NAME)

$$\bigcap_{C1}^{O}\bigcap_{NH}^{F}\bigcap_{NH}^{O}\bigcap_{F}^{Pr-n}$$

RN 122815-80-9 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)ethylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-81-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)ethylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-82-1 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)-2-propenylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-83-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)-2-propenylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-84-3 CAPLUS

CN Benzamide, N-[[[4-[[[(3,4-dichlorophenyl)-2-propenylamino]carbonyl]amino]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 122815-85-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(3,4-dichlorophenyl)-2-propenylamino]carbonyl]amino]-2-fluorophenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122815-86-5 CAPLUS

CN Benzamide, N-[[[4-[[(4-chlorophenyl)ethylamino]carbonyl]amino]-2-fluoro-5-methylphenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 122815-87-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)ethylamino]carbonyl]amino]-2-fluoro-5-methylphenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \circ & \circ & \text{F} \\ & \downarrow & \text{NH} & \downarrow & \text{NH} \\ & & \downarrow & \text{NH} & \downarrow & \text{NH} \\ & & & \text{C1} \\ \end{array}$$

RN 122815-88-7 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-2-fluoro-5-methylphenyl]amino]carbonyl]-2,6-difluoro-(9CI) (CA INDEX NAME)

RN 122815-89-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[(4-chlorophenyl)propylamino]carbonyl]amino]-2-fluoro-5-methylphenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 122829-04-3 CAPLUS

CN Benzamide, N-[[[2,5-difluoro-4-[[(phenylpropylamino)carbonyl]amino]phenyl] amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

- L5 ANSWER 10 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1988:160301 CAPLUS Full-text
- DN 108:160301
- TI Studies on the transition metal thiocyanate complexes with thioureas containing sulfur-sulfur and oxygen-sulfur-sulfur-oxygen donor sequences
- AU Tembe, G. L.; Murty, A. S. R.
- CS Dep. Chem., Karnatak Univ., Dharwad, 580 003, India
- SO Current Science (1987), 56(24), 1277-9 CODEN: CUSCAM; ISSN: 0011-3891
- DT Journal
- LA English
- AB ML(SCN)2 [M = Co, Ni, L = BzNHC(S)NH(CH2)2NHC(S)NHBz, o-C6H4(NHC(S)NHPh)2; m = Ni, L = o- and p-C6H4(NHC(S)NHBz)2] were prepared The complexes were characterized by molar conductivity and magnetic moment data, IR and electronic spectra and thermal anal. The ligands coordinate through the S atoms. Ligand field parameters were calculated The Ni complexes are octahedral and the Co complexes are 4 coordinate.
- RN 113804-06-1 CAPLUS
- CN Nickel, [N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis[benzamide]-S]bis(thiocyanato-S)-(9CI) (CA INDEX NAME)

RN 113804-07-2 CAPLUS

CN Nickel, [N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[benzamide]-S]bis(thiocyanato-S)- (9CI) (CA INDEX NAME)

RN 113804-09-4 CAPLUS

CN Cobalt, [N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis[benzamide]-S]bis(thiocyanato-S)- (9CI) (CA INDEX NAME)

L5 ANSWER 11 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1987:42838 CAPLUS Full-text

DN 106:42838

TI Binucleating bis-N-acylthioureas - ligands in trimetallamacrocycles and polynuclear metal chelates

AU Koehler, R.; Kirmse, R.; Richter, R.; Sieler, J.; Hoyer, E.; Beyer, L.

CS Sekt. Chem., Karl-Marx-Univ., Leipzig, Fed. Rep. Ger.

SO Zeitschrift fuer Anorganische und Allgemeine Chemie (1986), 537, 133-44
CODEN: ZAACAB; ISSN: 0044-2313

DT Journal

LA German

AB By sym. linking of 2 bidentate N-acylthioureas 2 types of quadridentate bis-N-acylthioureas are available which act, after di-deprotonation as bis-bidentate S, O ligands towards polyvalent metal ions. They can form oligomeric or polymeric, cyclic or chain chelates. With 1,1,1',1'-tetraalkyl-3,3'-terephthaloylbisthioureas (H2L) oligomeric triangulo-trimetallamacrocycles Ni3L3 and Cu3L3 were obtained. They contain perimetric 27-membered rings, counting the internal oxygens, or 39-membered rings with the external S atoms on the other hand, i.e. equal chalcogen atoms are in cis-positions within each chelate unit around the 3 metal ions. The trimetallamacrocyclic structure was

10/616,959

proved by x-ray crystal and mol. structure anal. of Ni3L3 (alkyl = Et) or EPR of the corresponding Cu3L3. Diamine-linked bis-N-acylthioureas form insol. 1:1 polymeric chelates.

IT 104359-19-5P 104359-20-8P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 104359-19-5 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis[(methylimino)carbonothioyl]]bis- (9CI) (CA INDEX NAME)

RN 104359-20-8 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis[(ethylimino)carbonothioyl]]bis- (9CI) (CA INDEX NAME)

L5 ANSWER-12 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN -

AN 1987:18148 CAPLUS Full-text

DN 106:18148

TI N,N'-disubstituted bisacylthiourea derivatives

IN Beyer, Lothar; Koehler, Ronald; Hoyer, Eberhard; Hartung, Juergen

PA Karl-Marx-Universitaet Leipzig, Ger. Dem. Rep.

SO Ger. (East), 11 pp.

CODEN: GEXXA8

DT Patent

LA German

FAN CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI DD 229400 PRAI DD 1984-270354 GI	A1	19851106 19841206	DD 1984-270354	19841206 <

The title compds. [RCONHC(S)NR1]2Z [I; R = (un)substituted Ph; R1 = alkyl, aryl; Z = (un)substituted arylene, (CH2)n; n = 2-18] and II [R as above; X, X1 = (CH2)2, CH:CH] are prepared as chelating agents. Thus, 6.5 g BzNCS (preparation given) was added to a solution of 2.6 g N,N'-dimethyl-p-phenylenediamine and 1 g Et3N in 30 mL acetone, to give I (R = Ph, R1 = Me, Z = p-C6H4) (III). III (5 mmol) in 80 mL DMF was added to 1.25 g Ni(OAc)2.4H2O in 150 mL DMF, to give a polymeric III.Ni complex.

IT 104359-19-5P 104359-20-8P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of, as chelating agent)

RN 104359-19-5 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis[(methylimino)carbonothioyl]]bis- (9CI) (CA INDEX NAME)

RN 104359-20-8 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis[(ethylimino)carbonothioyl]]bis- (9CI) (CA INDEX NAME)

L5 ANSWER 13 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1985:422429 CAPLUS Full-text

DN 103:22429

TI Synthesis and spectroscopic properties of some new N,N'-disubstituted thioureas of potential biological interest

AU Sarkis, George Y.; Faisal, Essam D.

CS Coll. Sci., Univ. Baghdad, Baghdad, Iraq

SO Journal of Heterocyclic Chemistry (1985), 22(1), 137-40 CODEN: JHTCAD; ISSN: 0022-152X

DT Journal

LA English

OS CASREACT 103:22429

AB Thirty-six N,N'-disubstituted thioureas RNHCSNHR1 [R = Bz, Ph, 4-FC6H4; R1 = (un)substituted Ph, pyridyl, 4-quinolyl] were synthesized by the reaction of RNCS with R1NH2. The UV, IR and NMR spectral data are presented and discussed.

IT 70110-39-3P 87874-16-6P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 70110-39-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

RN 87874-16-6 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

L5 ANSWER 14 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1984:630162 CAPLUS Full-text

DN 101:230162

TI Benzoylurea compounds for pesticidal and pharmaceutical use

IN Brouwer, Marius S.; Grosscurt, Arnoldus C.

PA Duphar International Research B. V., Neth.

SO Eur. Pat. Appl., 31 pp. CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

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	PAT	ENT	NO.			KINI)	DAT	3	Α	PP	LICATION	NO.	DATE	
							•			-				 	
ΡI	EP	1167	129			A2		1984	40829	E	P	1983-201	862	19831230	<
	ΕP	1167	129			A3		1984	40926						
	ΕP	1167	129			B1		1988	31012						
		R:	AT,	BE,	CH,	DE,	FR,	GB,	, IT,	LI,	LU	, NL, SE			
	ΑT	3786	59			E		1988	31015	А	T	1983-201	862	19831230	<
	AU	8423	3614			A1		1984	40726	А	U	1984-236	14	19840119	<
	AU	5622	260			В2		1987	70604						
	BR	8400	234			Α		1984	40828	В	R	1984-234		19840119	<
	ZA	8400	1422			Α		1984	40926	Z	Α	1984-422		19840119	<
	US	4665	235			Α		1987	70512	U	S	1984-572	143	19840119	<
	CA	1247	7644			A1		1988	31227	С	Ά	1984-445	614	19840119	<

10/616,959

	DK	8400268	Α	19840725	DK	1984-268	19840120	<
	DK	159923	В	19901231				
	DK	159923	С	19910521				
	DD	219101	A5	19850227	DD	1984-259516	19840120	<
	ES	529033	A1	19850316	ES	1984-529033	19840120	<
	PL	139504	B1	19870131	PL	1984-245840	19840120	<
	HU	35477	0	19850729	HU	1984-263	19840123	<
	HU	193668	В	19871130				
	IL	70747	A1	19861130	IL	1984-70747	19840123	<
	JP	59176242	A2	19841005	JP	1984-9592	19840124	<
	JP	04014660	B4	19920313				
	CS	242896	B2	19860515	CS	1984-527	19840124	<
	SU	1375125	A3	19880215	SU	1984-3751717	19840618	<
	US	4710516	Α	19871201	US	1986-932296	19861119	<
PRAI	NL	1983-238	Α	19830124		•		
	EΡ	1983-201862	Α	19831230				
	US	1984-572143	A2	19840119				
GI								

CONHCONH

R2

CONHCONH

R3

NHCO)
$$n = XR^5$$

R4

About 74 title compds. I (R1 = halo; R2 = H, halo; R3 = H, or 1-2 substituents selected from Cl, Me, CF3; R4 = H or 1-3 substituents selected from halo, alkyl, alkoxy, haloalkyl, haloalkoxy; X = N, CH; n = 0, 1; R5 = H, C1-6 alkyl, C2-6 alkenyl, C3-6 cycloalkyl; if n = 0, and R5 = H, then R3 = H), insecticides, acaricides, and antitumor agents, were prepared E.g., treating 0.90 g 2,6-F2C6H3CONCO with 1.27 g H2NC6H4NPrC6H4Cl-4 in Et2O at room temperature gave 1.50 g N-(2,6- difluorobenzoyl)-N'-[4-[N-(4-chlorophenyl)-N-propylamino]phenyl]urea (II). At 1 mg/L, II gave 90-91% mortality of larvae of Pieris brassicae.

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      TT
      93275-07-1P
      93275-08-2P
      93275-09-3P

      93275-35-5P
      93275-36-6P
      93275-37-7P

      93275-38-8P
      93275-39-9P
      93275-40-2P

      93275-41-3P
      93275-42-4P
      93275-43-5P

      93275-44-6P
      93275-45-7P
      93275-46-8P

      93275-50-4P
      93275-51-5P
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      93275-53-7P
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      93275-55-9P

      93275-56-0P
      93275-57-1P
      93275-58-2P

      93275-62-8P
      93275-63-9P
      93275-64-0P

      93275-72-0P
      93275-73-1P
      93275-74-2P

      93442-91-2P
      93442-91-2P
```

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation, pesticidal activity, and antitumor activity of)

RN 93275-07-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[(4-chlorophenyl)(1-

methylethyl)amino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-08-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]p henyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-09-3 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-35-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)ethylamino]carbonyl]amino]ph enyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-36-6 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-chlorophenyl)amino]carbonyl]amino]phenyl]amin o]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

RN 93275-37-7 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-chlorophenyl)amino]carbonyl]amino]phenyl]amin o]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-38-8 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-chlorophenyl)amino]carbonyl]amino]-3-chlorophenyl]amino]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

RN 93275-39-9 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-chlorophenyl)amino]carbonyl]amino]-3-chlorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-40-2 CAPLUS

CN Benzamide, N-[[[4-[[[butyl[4-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenyl]amino]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

RN 93275-41-3 CAPLUS

CN Benzamide, N-[[[4-[[[butyl[4-(trifluoromethyl)phenyl]amino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-42-4 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-methylphenyl)amino]carbonyl]amino]phenyl]amin o]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

RN 93275-43-5 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-methylphenyl)amino]carbonyl]amino]phenyl]amin o]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-44-6 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-methylphenyl)amino]carbonyl]amino]-3-chlorophenyl]amino]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ & \text{NH} \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline & \circ & \circ & \circ & \circ & \circ \\ \hline &$$

RN 93275-45-7 CAPLUS

CN Benzamide, N-[[[4-[[[butyl(4-methylphenyl)amino]carbonyl]amino]-3-chlorophenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-46-8 CAPLUS

CN Benzamide, N-[[[4-[[[butyl[4-(1,1,2,2-tetrafluoroethoxy)phenyl]amino]carbonyl]amino]phenyl]amino]carbonyl]-2-chloro- (9CI) (CA INDEX NAME)

RN 93275-47-9 CAPLUS

CN Benzamide, N-[[[4-[[[butyl[4-(1,1,2,2-tetrafluoroethoxy)phenyl]amino]carbo nyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-48-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[propyl[4-(1,1,2,2-tetrafluoroethoxy)phenyl]amino]carbonyl]amino]phenyl]amino]carbonyl](9CI) (CA INDEX NAME)

RN 93275-49-1 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[4-[[[propyl[4-(1,1,2,2-tetrafluoroethoxy)phenyl]amino]carbonyl]amino]phenyl]amino]carbonyl](9CI) (CA INDEX NAME)

RN 93275-50-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)(2-methylpropyl)amino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-51-5 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)(2-methylpropyl)amino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-52-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)hexylamino]carbonyl]amino]ph enyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

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 & & \\$$

RN 93275-53-7 CAPLUS

CN Benzamide, N-[[[4-[[(4-chlorophenyl)hexylamino]carbonyl]amino]phenyl]amin o]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-54-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)pentylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-55-9 CAPLUS

CN Benzamide, N-[[[4-[[(4-chlorophenyl)pentylamino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-56-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(2,6-dichlorophenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-57-1 CAPLUS

CN Benzamide, N-[[[4-[[[(2,6-dichlorophenyl)propylamino]carbonyl]amino]phenyl amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-58-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(3,4-dimethylphenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-59-3 CAPLUS

CN Benzamide, N-[[[4-[[[(3,4-dimethylphenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-60-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-fluorophenyl)propylamino]carbonyl]amino]p henyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-61-7 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[4-[[(4-fluorophenyl)propylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-62-8 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]p henyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-63-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-3-methylphenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-64-0 CAPLUS

CN Benzamide, N-[[[4-[[(4-chlorophenyl)propylamino]carbonyl]amino]-3-methylphenyl]amino]carbonyl]-2,6-difluoro-(9CI) (CA INDEX NAME)

RN 93275-65-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)-2-propenylamino]carbonyl]amino]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

$$\bigcap_{C1}^{O}\bigcap_{NH}^{O}\bigcap_{NH}^{C-NH}\bigcap_{NH}^{O}\bigcap_{NH}^{C+2-CH}\bigcap_{C1}^{C+2}$$

RN 93275-66-2 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)-2-propenylamino]carbonyl]amino]pheny l]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} F & O & O & CH_2-CH = CH_2 \\ \hline C & NH & C & NH & C \\ \hline \end{array}$$

RN 93275-71-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-3,5-dimethylphenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-72-0 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-3,5-dimethylphenyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

RN 93275-73-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-3-(trifluoromethyl)phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 93275-74-2 CAPLUS

CN Benzamide, N-[[[4-[[[(4-chlorophenyl)propylamino]carbonyl]amino]-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro-(9CI) (CA INDEX NAME)

RN 93442-91-2 CAPLUS

CN Benzamide, N-[[[4-[[(4-chlorophenyl)(1-methylethyl)amino]carbonyl]amino]p henyl]amino]carbonyl]-2,6-difluoro- (9CI) (CA INDEX NAME)

L5 ANSWER 15 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1984:6377 CAPLUS Full-text

DN 100:6377

TI Reactions of carbonyl isothiocyanates with nucleophilic bifunctional reagents

AU Uher, Michal; Berkes, Dusan; Lesko, Jan; Floch, Lubomir

CS Dep. Org. Chem., Slovak Inst. Technol., Bratislava, 812 37, Czech.

SO Collection of Czechoslovak Chemical Communications (1983), 48(6), 1651-8
CODEN: CCCCAK; ISSN: 0366-547X

DT Journal

LA English

OS CASREACT 100:6377

GΙ

AB Acyl isothiocyanates RCONCS (R = Me, Cl3C, Ph, 2-furanyl) condensed with I (X = CH, Z = O, S, NH; X = N, Z = NH) and II to give acylthioureas. Those derived from I (X = CH, Z = S, NH) were cyclized with elimination of H2S to give III.

IT 87874-16-6P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 87874-16-6 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

L5 ANSWER 16 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1983:487771 CAPLUS Full-text

DN 99:87771

TI Studies on the alkoxybenzoic acid series. V. 3,4,5-Trimethoxybenzoyl thioureides

AU Missir, A.; Zolta, V.; Soare, Jana; Chirita, Ileana; Petrea, I.; Stan, A.

CS Lab. Chim. Farm., Fac. Farm., Bucharest, Rom.

SO Farmacia (Bucharest, Romania) (1982), 30(4), 225-30 CODEN: FRMBAZ; ISSN: 0014-8237

DT Journal

LA Romanian

OS CASREACT 99:87771

GΙ

AB Bis-thioureas I [Z = phenylene, methylphenylene, (CH2)n (n = 2,3,4,5,6)] and benzoylthioureas II [R = 3,4,5-(MeO)3C6H2CONHCS, Ph] were prepared Thus, 3,4,5-(MeO)3C6H2COC1 was treated with NH4SCN in Me2CO, the mixture was heated, o-phenylenediamine in Me2CO was added, and the mixture was refluxed to give I (Z = o-phenylene).

IT 82925-65-3P 82925-69-7P 82934-52-9P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 82925-65-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

RN 82925-69-7 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis[3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

RN 82934-52-9 CAPLUS

CN Benzamide, N,N'-[(2-methyl-1,4-phenylene)bis(iminocarbonothioyl)]bis[3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

- L5 ANSWER 17 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1982:555973 CAPLUS Full-text
- DN 97:155973
- TI Pharmacodynamic study of some new 3,4,5-trimethoxybenzoic acid thioureides. Part VI
- AU Cristea, Elena; Missir, A.; Chirita, Ileana; Dan, G.; Georgescu, C.
- CS Discip. Farmacodin., Fac. Farm., Bucharest, Rom.
- SO Farmacia (Bucharest, Romania) (1982), 30(1), 41-8 CODEN: FRMBAZ; ISSN: 0014-8237
- DT Journal
- LA Romanian

GI

The pharmacol. of 11 title compds. [I(Z = (CH2)n, n = 2-6, etc.); II (R = 4-Ph-piperazin-1-yl or 2,6-Br2C6H3NH) and III [82925-64-2]] was studied. Among the central nervous system depressing substance were I (Z = p-C6H4) [82925-65-3], I [Z = (CH2)3] [82925-66-4], I [Z = (CH2)5] [82925-67-5], II (R = 4-Ph-piperazin-1-yl, and III. Compds. blocking intestinal motility included I (Z = o-C6H4) [82925-69-7], I (Z = p-C6H4), I [Z = (CH2)4] [82925-70-0], and I (Z = 2-Me-1,4-C6H3. The compds. had anticholesteremic and antihyperglycemic activities. None of the compds. had greater activity than compds. of the same class previously tested.

IT 82925-65-3 82925-69-7 82934-52-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pharmacol. of)

RN 82925-65-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{MeO} \\ \text{MeO$$

RN 82925-69-7 CAPLUS

CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonothioyl)]bis[3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

RN 82934-52-9 CAPLUS

CN Benzamide, N,N'-[(2-methyl-1,4-phenylene)bis(iminocarbonothioyl)]bis[3,4,5-trimethoxy-(9CI) (CA INDEX NAME)

L5 ANSWER 18 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1982:227948 CAPLUS Full-text

DN 96:227948

TI Complexes of p,p'-bis(benzoylthioureido)benzene with copper(II), nickel(II) and cobalt(II) salts and their biological activity

AU Satpathy, K. C.; Mishra, H. P.; Patel, B. N.

CS P. G. Dep. Chem., Sambalpur Univ., Burla, 768 017, India

SO Journal of the Indian Chemical Society (1982), 59(1), 40-2 CODEN: JICSAH; ISSN: 0019-4522

DT Journal

LA English

AB MLX2 (M = Cu, Ni, Co; L = BzNHC(S)NHC6H4NHC(S)NHBz-p, X = Cl, Br, NO3, ClO4) were prepared and characterized on the basis of IR spectral, electronic spectra and magnetic susceptibility measurements. IR spectra manifest the coordinates of the ligand to the metal ion through carbonyl O and thiocarbonyl S atoms. The complexes possess octahedral stereochem. as inferred from electronic spectral data and magnetic moment values. Fungicidal screening of the complexes shows them to be antifungal against Aspergellus niger, Fusarium oxysporium and Helminthosporium oryzae.

IT 70110-39-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and fungicidal activity of)

RN 70110-39-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

L5 ANSWER 19 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1982:85276 CAPLUS Full-text

DN 96:85276

TI 1-(2-Aminophenyl)-3-phosphonoureas and thio analogs

IN Weir, W. David; Kilbourn, Edward E.

PA Rohm and Haas Co., USA

SO U.S., 14 pp. Division of U.S. 4,183,921.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 8

	PATENT NO.		KIND	DATE	API	PLICATION NO.	DATE		
PI	US 427	76290	Α	19810630	US	1979-73249	19790906	<	
	JP 490	048641	A2 .	19740511	JP	1973-60196	19730529	<	
	CA 101	17351	A1	19770913	CA	1973-172986	19730601	<	
	AT 740	09010	Α	19770715	AT.	1974-9010	19741111	<	
	AT 342	2362	В	19780328					
	ES 441	1218	A1	19770701	ES	1975-441218	19750916	<	
	ES 441	1218	A5	19770708					
	US 407	76809	Α	19780228	US	1975-625998	19751028	<	
	US 418	33921	Α	19800115	US	1977-842645	19771017	<	
PRAI	US 197	72-263378	A2	19720605					
	US 197	73-354629	A2	19730425					
	US 197	75-625998	A3	19751028					
	US 197	77-842645	A3	19771017					
	US 197	72-259423	Α	19720605					
	AT 197	73-4588	Α	19730525					
•	ES 197	73-415199	A3	19730525					

OS CASREACT 96:85276; MARPAT 96:85276

Ureas and analogs RZNHC(X)NR1P(X1)R2R3 (R = NH2, heteroarom. acylamino, substituted ureido or thioureido, substituted 3-phosphonoureido and thio analogs; Z = phenylene, naphthalenediyl, anthracenediyl, phenanthrenediyl, diazopinediyl, pyridinediyl, pyrimidinediyl, furandiyl; X = O, S; R1 = H, hydrocarbyl; X1 = O, S; R2 = hydrocarbyl, hydrocarbyloxy, substituted amino, hydrocarbylthio; R3 = Hydrocarbyloxy, substituted amino, hydrocarbylthio), useful as anthelmintics (no data), were prepared Thus, o-phenylenediamine was treated with (EtO)2PONCS to give 2-H2NC6H4NHC(S)NHP(O)(OEt)2.

IT 52406-05-0P 52406-06-1P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 52406-05-0 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]carbonyl]amino]phenyl] amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-06-1 CAPLUS

CN Phosphoramidic acid, [[[2-[[(4-nitrobenzoyl)amino]carbonyl]amino]phenyl]a mino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 20 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1979:187379 CAPLUS Full-text

DN 90:187379

TI Synthesis of polyacylthioureas by polyaddition of isophthaloyldiisothiocyanate with diamines

AU Shimano, Yasuo; Sasaki, Shoichi

CS Dep. Ind. Chem., Hachinohe Tech. Coll.; Hachinohe, Japan

SO Kobunshi Ronbunshu (1979), 36(2), 81-8

CODEN: KBRBA3; ISSN: 0386-2186

DT Journal

LA Japanese

AB Isophthaloyl diisothiocyanate (I) is polymd. with arom. diamines in amide solns. to give polymers having reduced viscosity ≤1.39 dL/g (30°, 0.5 g/dL in Me2NAc containing 5% LiCl), or I is polymerized with aliphatic diamines by interfacial methods using aromatic solvents to give polymers having reduced viscosity up to 1.21 dL/g. Interfacial polymerization of I with aromatic diamines and solution polymerization of I in amide solvents with aliphatic diamines does not give high-mol. weight polymers. The poly(acylthioureas) lose 5% weight in N or air at 210-20°.

IT 70113-14-3P 70113-15-4P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and properties of, solvent effect on)

RN 70113-14-3 CAPLUS

CN Poly(iminocarbonothioylimino-1,4-phenyleneiminocarbonothioyliminocarbonyl-1,3-phenylenecarbonyl) (9CI) (CA INDEX NAME)

Poly(iminocarbonothioylimino-1,3-phenyleneiminocarbonothioyliminocarbonyl-CN 1,3-phenylenecarbonyl) (9CI) (CA INDEX NAME)

$$\left[\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

IT 70110-39-3P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN70110-39-3 CAPLUS

Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CI) CN

L5 ANSWER 21 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

1978:508754 CAPLUS Full-text AN

DN 89:108754

TI · Anthelmintic phosphorothioureidoarylenethioureas

IN Owen, Ronald P.; Miller, George A.; Schneider, Charles M.

PA Rohm and Haas Co., USA

SO U.S., 14 pp.

CODEN: USXXAM

DTPatent

LA English

FAN.CNT 8

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	US 4086336	 A	19780425	us 1976-717411	 19760824 <
	ZA 7303432	A	19740626	ZA 1973-3432	19730522 <
	BR 7303866	A0	19731220	BR 1973-3866	19730525 <
	JP 49048640	A2	19740511	JP 1973-60195	19730529 <
	AT 7409010	Α	19770715	AT 1974-9010	19741111 <
	AT 342362	В	19780328		
	US 4170648	Α	19791009	US 1978-876779	19780210 <
PRAI	US 1972-259423	A2	19720526		
	US 1973-354630	A2	19730425		
	US 1972-263378	Α	19720605		
	AT 1973-4588	Α	19730525		
	US 1976-717411	A3	19760824		
os	MARPAT 89:108754				

MARPAT 89:108754

The title compds., R3NHCSNHXNHCSNRP(O)R1R2 (I, X = arylene optionallysubstituted with halo, cyano, thiocyano, carboxy, nitro, amino, etc.; R = H, C1-10 alkyl, C1-10 haloalkyl, C3-6 cycloalkyl, C2-11 alkoxyalkyl, C1-10 cyanoalkyl, C3-6 alkenyl, C3-6 haloalkenyl, C3-6 alkynyl, C3-6 haloalkynyl,

C7-11 aralkyl, C6-10 aryl; R1, R2 = R4, OR4, NR42, SR4, R4 = C1-18 alkyl; R3 = MeSO2, Ac, EtCO, ClCH2CO, Bz, P(O)R5R6, R5, R6 = OR4, NR42, SR4) were prepared by the reaction of o-(H2N)2X with R1R2P(O)NCS to give o-H2NXNHCSNHP(O)R1R2, which, with R3NCS, gave I. Thus, o-(H2N)2C6H4 and (EtO)2P(O)NCS gave o-H2NC6H4NHCSNHP(O)(OEt)2, which, with MeSO2NCS, gave o-MeSO2NHCSNHC6H4NHCSNHP(O)(OEt)2. Among the 20 I similarly prepared were o-(Me2N)2P(O)NHCSNHC6H4NHCSNHP(O)(OEt)(SMe), o-BZNHCSNHC6H4NHCSNHP(O)(OEt)2, and o-(Me2CHS)(EtO)P(O)NHCSNHC6H4NHCSNHP(O)(OEt)(SCHMe2). The preferred dose of I for roundworms in animals is 12-100 mg/kg.

IT 52406-18-5P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 52406-18-5 CAPLUS

CN Phosphoramidic acid, [[[2-[[(benzoylamino)thioxomethyl]amino]phenyl]amino] thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 22 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1978:424007 CAPLUS Full-text

DN 89:24007

TI Phosphonothioureides

IN Weir, William David

PA Rohm and Haas Co., USA

SO Patentschrift (Switz.), 7 pp.

CODEN: SWXXAS

DT Patent

LA German

FAN.CNT 2

L MIA .	PATENT NO.		DATE	APPLICATION NO.	DATE
PI	CH 596228	A	19780315	CH 1973-14623	19731016 <
	US 3845176	Α	19741029	US 1972-298683	19721018 <
	FR 2306700	A2	19761105	FR 1973-36312	19731011 <
	FR 2306700	B2	19790126		
	BE 806083	A4	19740416	BE 1973-136693	19731015 <
	ZA 7307995	Α	19741127	ZA 1973-7995	19731015 <
	DD 109223	W	19741020	DD 1973-174091	19731016 <
	AU 7361459	A1	19750417	AU 1973-61459	19731016 <
	JP 54007787	B4	19790410	JP 1973-116249	19731016 <
	SE 415355	В	19800929	SE 1973-14069	19731016 <
	SE 415355	С	19810122		
	GB 1444103	Α	19760728	GB 1973-48353	19731017 <
	ни 172069	P	19780528	HU 1973-RO754	19731017 <
	NL 7314380	A	19740422	NL 1973-14380	19731018 <
	AT 7308868	Α	19760315	AT 1973-8868	19731018 <
	AT 333305	В	19761110		•
	ES 419749	A1	19760316	ES 1973-419749	19731018 <
	PL 101308	P	19781230	PL 1973-165936	19731018 <
	IL 43491	A1	19780310	IL 1973-43491	19731026 <

IN 139438 A 19760619 IN 1974-CA403 19740226 <-PRAI US 1972-298683 A 19721018
BE 1973-800041 A 19730525

GΙ

AB Twenty-two phosphonothioureides I (R = H or the same or different alkyl, alkoxy, halo; R1 = alkylsulfonyl, PhSO2, substituted phenylsulfonyl, alkanoyl, Bz, substituted benzoyl; R2 = alkyl, alkoxyalkyl, haloalkyl, Ph, substituted phenyl; X = O or S; n = 1-4) were prepared by treating RnC6H4-n(NH2)2 (NH2 groups ortho to one another) successively with SCNP(O) (OR2)2 and XCNR1.

IT 52406-05-0P 52406-06-1P 52406-18-5P

52406-19-6P 52406-20-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 52406-05-0 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]carbonyl]amino]phenyl] amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-06-1 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-nitrobenzoyl)amino]carbonyl]amino]phenyl]a mino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-18-5 CAPLUS

CN Phosphoramidic acid, [[[2-[[(benzoylamino)thioxomethyl]amino]phenyl]amino] thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-19-6 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]thioxomethyl]amino]phe nyl]amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-20-9 CAPLUS

CN Phosphoramidic acid, [thioxo[[2-[[thioxo[[4-(trichloromethyl)benzoyl]amino]methyl]amino]methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 23 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1978:406131 CAPLUS Full-text

DN 89:6131

TI 1-(2-Ureidophenyl)-3-phosphonoureas

IN Weir, W. David; Kilbourn, Edward E.

PA Rohm and Haas Co., USA

SO U.S., 17 pp.

CODEN: USXXAM

DT Patent

LA English

FAN. CNT 8

EMM.	CNIO							
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE			
PI	US 4076809	Α	19780228	US 1975-625998	19751028 <			
	US 4183921	Α	19800115	US 1977-842645	19771017 <			
	US 4234575	Α	19801118	US 1979-7881 .	19790131 <			
	US 4276290	Α	19810630	US 1979-73249	19790906 <			
PRAI	US 1972-263378	A2	19720605					
	US 1973-354629	A2	19730425					
	US 1975-625998	A3	19751028					
	US 1977-842645	A3	19771017					
GI								

AB ZANHC(X)NRP(X1)YY1 (Z = NH2, acylamino, a ureido group, and thioureido group; A = substituted or unsubstituted divalent arylene; X = O, S; R = H, alkyl, haloalkyl, cycloalkyl, alkoxyalkyl, cyanoalkyl, alkenyl, haloalkenyl, alkynyl, haloalkynyl, aralkyl, substituted or unsubstituted aryl; X1 = O, S; Y = alkyl, alkenyl, aryl, alkoxy, alkenyloxy, aryloxy, alkylthio, alkenylthio, disubstituted amino; Y1 = alkoxy, alkenyloxy, aryloxy, alkylthio, alkenylthio, arylthio, disubstituted amino), which are useful as anthelmintics (no data), were prepared Thus, o-phenylenediamines was acylated by (EtO)2P(O)NCS to give I (R = H) which with PhSO2NCO yielded I (R = CONHSO2Ph).

IT 52406-05-0P 52406-06-1P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 52406-05-0 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]carbonyl]amino]phenyl] amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-06-1 CAPLUS

CN Phosphoramidic acid, [[[2-[[(4-nitrobenzoyl)amino]carbonyl]amino]phenyl]a mino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 24 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1976:483225 CAPLUS Full-text

DN 85:83225

TI Anthelmintics containing phosphonothioureides

PA Rohm and Haas Co., USA

SO Jpn. Kokai Tokkyo Koho, 9 pp. CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE			
		-						
ΡI	JP 50050341	A2	19750506	JP 1973-98162	19730831 <			
PRAI	JP 1973-98162	Α	19730831					
GI								

AB Anthelmintics for domestic animals and humans contain phosphonothioureides I and II (R2 = lower alkyl, substituted or unsubstituted aryl; R1 = H, lower alkyl, lower alkoxy, or halogen; R3, R4 = alkyl or substituted or unsubstituted aryl). For example, I (R1 = H, R2 = R3 = Et) (IV) [52406-12-9] was prepared from III [52405-98-8] and EtSO2NCS [52405-94-4]. A tasty tablet preparation contained active ingredient (such as IV) 110, fish powder 1027, bovine liver powder 1027, soybean powder 97 and sucrose 239 mg.

IT 52406-18-5P 52406-19-6P 52406-20-9P

RL: PREP (Preparation)

(preparation of, for anthelmintic pharmaceuticals)

RN 52406-18-5 CAPLUS

CN Phosphoramidic acid, [[[2-[[(benzoylamino)thioxomethyl]amino]phenyl]amino] thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-19-6 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]thioxomethyl]amino]phe nyl]amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-20-9 CAPLUS

CN Phosphoramidic acid, [thioxo[[2-[[thioxo[[4-(trichloromethyl)benzoyl]amino]methyl]amino]methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 25 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1976:468293 CAPLUS Full-text

DN 85:68293

TI Phosphonothioureide anthelmintics

PA Rohm and Haas Co., USA

SO Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 8

212110111				
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI JP 50035134	A2	19750403	JP 1974-45884	19740423 <
PRAI US 1973-354630	Α	19730425	•	
GI .				

- AB Anthelmintics for human and domestic animals contain phosphonothioureids(I) (R = lower alkyl, substituted or unsubstituted aryl; Rl = H, lower alkyl, lower alkoxy or halogen; R2 = SO2R3 or COR4 with R3 and R4 = alkyl or substituted or unsubstituted aryl). E.g., I(R = Et, Rl = H, R2 = SO2Et) (II) [52406-12-9] was prepared from III [52405-98-8] and EtSO2NCS [52405-94-4]. A tablet preparation contained active ingredient (such as II) 220, lactose 53.23, MgAl silicate 2.24, starch 3.13 Ca stearate 0.65 and microcryst. cellulose 35.75.
 - 52406-18-5P 52406-19-6P 52406-20-9P

RL: PREP (Preparation)

(preparation of, for anthelmintic pharmaceuticals)

RN 52406-18-5 CAPLUS

IT

CN Phosphoramidic acid, [[[2-[[(benzoylamino)thioxomethyl]amino]phenyl]amino] thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-19-6 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]thioxomethyl]amino]phe nyl]amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-20-9 CAPLUS

CN Phosphoramidic acid, [thioxo[[2-[[thioxo[[4-(trichloromethyl)benzoyl]amino]methyl]amino]methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 26 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1976:437249 CAPLUS Full-text

DN 85:37249

TI Phosphonoureide and phosphonothioureide anthelmintics for domestic animals

PA Rohm and Haas Co., USA

SO Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 8

	0111 .0			•	
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΙ	JP 49135952	A2	19741227	JP 1973-98163	19730831 <
PRAI	US 1973-354629	Α	19730425		
GT					

AB Anthelmintics for dosmetic animals contain phosphonoureides or phosphonothioureides (I) (X and Y = O or S; R = lower alkyl, lower alkoxyalkyl, aryl, halogenated lower alkyl, substituted or unsubstituted phenyl; R1 = H, lower alkyl, lower alkoxy or halogen; R3 = SO2R3 or COR4 with

R3 and R4 = alkyl or substituted or unsubstituted aryl). E.g. I(R = Et, R1 = H, R2 = SO2PH, X = S, Y = O) (II) [52406-00-5] was prepared from III [52405-98-8] and PhSO2NCO [2845-62-7]. A tablet preparation contained active ingredient (such as II) 220, lactose 53.23, MgAl silicate 2.24, starch 13.13, Ca stearate 0.65 and microcryst. cellulose 35.75 mg.

IT 52406-05-0P 52406-06-1P

RL: THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of, as anthelmintic)

RN 52406-05-0 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]carbonyl]amino]phenyl] amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-06-1 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-nitrobenzoyl)amino]carbonyl]amino]phenyl]a mino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 27 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1974:437407 CAPLUS Full-text

DN 81:37407

TI 1-(3-Disubstituted phosphonothioureido)-2-(3-substituted ureido- or thioureido)-benzene compounds

IN Weir, William D.

PA Rohm and Haas Co.

SO Ger. Offen., 24 pp. CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 2

T.T.TA	· CNI Z				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI ·	DE 2346241	A1	19740502	DE 1973-2346241	19730913 <
	US 3845176	Α	19741029	US 1972-298683	19721018 <
	FR 2306700	A2	19761105	FR 1973-36312	19731011 <
	FR 2306700	B2	19790126		
	BE 806083	A4	19740416	BE 1973-136693	19731015 <
	ZA 7307995	Α	19741127	ZA 1973-7995	19731015 <
	DD 109223	W	19741020	DD 1973-174091	19731016 <
	AU 7361459	A1	19750417	AU 1973-61459	19731016 <

JP	54007787	B4	19790410	JP	1973-116249	19731016	<
SE	415355	В	19800929	SE	1973-14069	19731016	<
SE	415355	С	19810122				
GB	1444103	Α	19760728	GB	1973-48353	19731017	<
HU	172069	P	19780528	HU	1973-R0754	19731017	<
NL	7314380	Α	19740422	NL	1973-14380	19731018	<
AT	7308868	Α	19760315	ΑT	1973-8868	19731018	<
AT	333305	В	19761110				
ES	419749	A1	19760316	ES	1973-419749	19731018	<
PL	101308	P	19781230	PL	1973-165936	19731018	<
IL	43491	A1	19780310	ΙL	1973-43491	19731026	<
IN	139438	Α	19760619	IN	1974-CA403	19740226	<
PRAI US	1972-298683	Α	19721018				
BE	1973-800041	Α	19730525				

GI For diagram(s), see printed CA Issue.

The urea derivs. I (R = Et, Me2CH, ClCH2CH2; Rl = H, Cl; R2 = e.g., 4-MeC6H4SO2, BuSO2, Ac, Bz; Z = O, S) were prepared in one reaction vessel by the reaction of ClP(O) (OR) 2 with a thiocyanate to give SCNP(O) (OR) 2, which reacted with 3,4-(H2N)2C6H3R, then with R2NCS or R2NCO to give I. Thus, ClP(O) (OEt)2 reacted with KSCN in MeOCH2CH2OMe, followed by addition of o-C6H4(NH2)2, then 4-MeC6H4SO2NCS to give I (R = Et, Rl = H, R2 = 4-MeC6H4SO2, Z = S). Twenty-two I were prepared

IT 52406-05-0P 52406-06-1P 52406-18-5P

52406-19-6P 52867-32-0P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 52406-05-0 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]carbonyl]amino]phenyl] amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-06-1 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-nitrobenzoyl)amino]carbonyl]amino]phenyl]a mino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-18-5 CAPLUS

CN Phosphoramidic acid, [[[2-[[(benzoylamino)thioxomethyl]amino]phenyl]amino] thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-19-6 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]thioxomethyl]amino]phe nyl]amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52867-32-0 CAPLUS

CN Phosphoramidic acid, [thioxo[[4-[[thioxo[[4-(trichloromethyl)benzoyl]amino]methyl]amino]methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 28 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1974:95531 CAPLUS Full-text

DN 80:95531

TI Anti-worm arylene compounds

IN Schneider, Charles M.; Owne, Ronald Parris; Miller, George Allen; Wier, William D.; Kilbourn, Edward E.

PA Rohm and Haas Co.

SO Ger. Offen., 32 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 8

L 1 H 1					
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	DE 2325036	A1	19740103	DE 1973-2325036	19730517 <
	DE 2325036	в2	19800313		
	DE 2325036	C3	19801106		
	ZA 7303432	Α	19740626	ZA 1973-3432	19730522 <
	NL 7307291	Α	19731207	NL 1973-7291	19730524 <
	SE 411758	В	19800204	SE 1973-7360	19730524 <
	SE 411758	С	19800522		•
	BE 800041	A1	19731126	BE 1973-131524	19730525 <
	BR 7303866	A0	19731220	BR 1973-3866	19730525 <

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FR 2187336
                           A1
                                 19740118
                                              FR 1973-19183
                                                                       19730525 <--
     FR 2187336
                           В1
                                 19770128
     DD 105441
                           С
                                 19740420
                                              DD 1973-171078
                                                                       19730525 <--
     AU 7356165
                           A1
                                 19741128
                                              AU 1973-56165
                                                                       19730525 <--
     AT 7304588
                           Α
                                 19750215
                                              AT 1973-4588
                                                                       19730525 <--
     AT 326145
                           В
                                 19751125
     CH 574451
                           Α
                                 19760415
                                              CH 1973-7569
                                                                       19730525 <--
     IN 139086
                           Α
                                 19760508
                                              IN 1973-CA1229
                                                                       19730525 <--
     ES 415199
                           A1
                                 19760601
                                              ES 1973-415199
                                                                       19730525 <--
     HU 169608
                           Р
                                 19761228
                                              HU 1973-R0733
                                                                       19730525 <--
     IL 42355
                           A1
                                 19771230
                                              IL 1973-42355
                                                                       19730525 <--
     CS 193022
                           Р
                                 19790917
                                              CS 1973-3805
                                                                       19730525 <--
     DK 142752
                           В
                                 19810112
                                              DK 1973-2903
                                                                       19730525 <--
     JP 49048640
                           A2
                                 19740511
                                              JP 1973-60195
                                                                       19730529 <--
     JP 49048641
                          · A2
                                 19740511
                                              JP 1973-60196
                                                                       19730529 <--
     GB 1389525
                           Α
                                 19750403
                                              GB 1973-26150
                                                                       19730601 <--
     SU 536755
                           D
                                 19761125
                                              SU 1973-1926076
                                                                      19730604 <--
     AT 7409010
                           Α
                                 19770715
                                              AT 1974-9010
                                                                      19741111 <--
     AT 342362
                                 19780328
                           В
                                              ES 1975-441218
     ES 441218
                           A1
                                 19770701
                                                                      19750916 <--
     ES 441218
                                 19770708
                           A5
PRAI US 1972-259423
                                 19720526
                           Α
     US 1972-263378
                           Α
                                 19720605
     AT 1973-4588
                           Α
                                 19730525
     ES 1973-415199
                           A3
                                 19730525
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GI For diagram(s), see printed CA Issue.

AB Urea derivs. (I; R = Et, C1CH2CH2, Me2CH; R1 = e.g., Ac, EtCO, 4-O2NC6H4CO, MeSO2, PhSO2, 4-MeC6H4SO2; Z = O,S) were prepared by the reaction of o-C6H4(NH2)2 with (RO)2P(:Z)NCS or (RO)2P(:Z)NCO followed by reaction with R1NCS or R1NCO. About 20 compds. were prepared, useful as nematocides in e.g., sheep and cattle.

IT 52406-05-0P 52406-06-1P 52406-18-5P

52406-19-6P 52406-20-9P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 52406-05-0 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]carbonyl]amino]phenyl] amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-06-1 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-nitrobenzoyl)amino]carbonyl]amino]phenyl]a mino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-18-5 CAPLUS

CN Phosphoramidic acid, [[[2-[[(benzoylamino)thioxomethyl]amino]phenyl]amino] thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-19-6 CAPLUS

CN Phosphoramidic acid, [[[2-[[[(4-methylbenzoyl)amino]thioxomethyl]amino]phe nyl]amino]thioxomethyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 52406-20-9 CAPLUS

CN Phosphoramidic acid, [thioxo[[2-[[thioxo[[4-(trichloromethyl)benzoyl]amino]methyl]amino]methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 29 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1972:564599 CAPLUS Full-text

DN 77:164599

TI Formation of 2-benzamidobenzimidazole. Its thermal stability

AU Depost, Gerard; Salle, Robert; Sillion, Bernard

CS Dep. Rech., Inst. Fr. Pet., Grenoble, Fr.

SO Comptes Rendus des Seances de l'Academie des Sciences, Serie C: Sciences

Chimiques (1972), 275(13), 697-700 CODEN: CHDCAQ; ISSN: 0567-6541

- DT Journal
- LA French
- GI For diagram(s), see printed CA Issue.
- 2-Benzamidobenzimidazole (I) was obtained in 17 yield by cyclizing o-H2NC6H4NHCONH-Bz with p-MeC6H4SO3H in boiling PhMe. In the absence of p-MeC6H4SO3H, o-H2NC6H4NHCONHBz decomposed to (H2N)2C6H4, o-(BzNHCONH)2C6H4, BzNH2, and 2-phenyl-benzimidazole. O-H2NC6H4NHCONHBz was prepared in 36 yield by treating BzNCO with o-(H2N)2C6H4 at room temperature Pyrolysis of I at 320° gave H2O, C6H6, PhCN, BzNH2, 2-phenylbenzimidazole, and the trimer II.
- IT 38870-79-0P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

- RN 38870-79-0 CAPLUS
- CN Benzamide, N,N'-[1,2-phenylenebis(iminocarbonyl)]bis- (9CI) (CA INDEX NAME)

- L5 ANSWER 30 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1971:449011 CAPLUS Full-text
- DN 75:49011
- TI New iodinated organic compounds. Iodinated derivatives of 1,2-dihydro-4H-3,1-benzoxazine-2,4-dione and 2,4(1H, 3H)-quinazolinedione
- AU Covello, Mario; Dini, Antonio; De Simone, Francesco
- CS Ist. Chim. Farm. Tossicol., Univ. Napoli, Naples, Italy
- SO Rendiconto dell'Accademia delle Scienze Fisiche e Matematiche, Naples (1969), 36, 61-6 CODEN: RASFAM; ISSN: 0370-3568
- DT Journal
- LA Italian
- GI For diagram(s), see printed CA Issue.
- AB The known 6,2-I(H2N)C6H3CO2H (I) refluxed 20 hr in ClCO2Et yielded 63% 5-iodo-2H-3,1-benzoxazine-2,4-(1H)-dione (II) (R = H, R1 = 5-I), m. 173.5° (MeOH-C6H6), converted by refluxing 2 hr in concentrated NH4OH to 39% 5-iodo-2,4-(1H, 3H)-quinazolinedione (III) (R = H, R1 = 5-I), m. 340°, also produced by heating I 30 min at 170-80° with urea. NH4SCN refluxed in Me2CO with addition of BzCl and the mixture treated with I in Me2CO, refluxed and the cooled solution poured into cold H2O gave 6,2-I(BzNHCSNH)C6H3CO2H (IV), m. 171-3°, converted by refluxing in N NaOH and acidification to 5-iodo-2-thio-(2,4(1H,3H)-quinazolinedione (V) (R = H, R1 = 5-I), m. 324-6° (decomposition).The known 3.5.2-ICl(NH2)C6H2CO2H was similarly transformed to give 46% II (R = 6-C1, R1 = 8-I), m. $176-8^{\circ}$; 62° III (R = 6-C1, R1 = 8-I), m. 310° (decomposition), 47% 3,5,2-ICl(BzNHCSNH)C6H2CO2H, m. $181-3^{\circ}$, and 80% V (R = 6-Cl, R1 = 8-I), m. $320-2^{\circ}$ (decomposition). Analogous procedures converted 3,5,2-IBr(H2N)C6H2CO2H into 88% II (R = 6-Br, R1 = 8-I), m. $155-7^{\circ}$; 43% III $(R = 6-Br, R1 = 8-I), m. 314-16^{\circ}; 71\% acid 3,5,2-IBr(BzNHCSNH)C6H2CO2H, m.$ $172-4^{\circ}$; and 84% V (R = 6-Br, R1 = 8-I), m. 303-5° (decomposition).
- IT 33115-22-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 33115-22-9 CAPLUS

CN Benzoic acid, 2,6-bis(3-benzoyl-2-thioureido)- (8CI) (CA INDEX NAME)

L5 ANSWER 31 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1966:84555 CAPLUS Full-text

DN 64:84555

OREF 64:15870g-h,15871a-h,15872a-b

TI Thioacyl isocyanates. III. Synthesis and properties of N-thiobenzoylureas

AU Goerdeler, Joachim; Schenk, Hainfried

CS Univ. Bonn, Germany

SO Chemische Berichte (1966), 99(3), 782-92

CODEN: CHBEAM; ISSN: 0009-2940

DT Journal

LA German

OS CASREACT 64:84555

GI For diagram(s), see printed CA Issue.

AB cf. CA 64, 5083d. Primary and secondary amines were added to PhCSNCO (I) to yield the corresponding PhCSNHCONRR' (II). PhCSNHCONH2 (III) was obtained by the selective saponification of II (R = Bz, R' = H) (IV). The adducts from hydrazines and amidines to I showed a strong tendency for cyclization. Phenylthiazolidine-4,5-dione (V) (5 g.) in 30 cc. dry methylcyclohexane decomposed thermally by the method described previously gave a solution of I; except where noted otherwise, this solution from 5 g. V was used in all runs with I as the starting material. I treated dropwise with 1.2 g. absolute EtOH yielded 3 g. deep yellow PhCSNHCO2Et, 63° (decomposition) (AcOEt-ligroine). with 1.92 g. BuNH2 in 5 cc. dry Et2O gave after chromatography on silica gel 0.7 g. PhCN, 1.5 g. PhCSNH2, 0.28 g. II (R = Bu, R' = H), m. 92° (1:15 CH2C12methylcyclohexane), and 2 g. brown, odoriferous oil. I with 2.23 g. piperidine in 25 cc. dry methylcyclohexane stirred 15 min. gave 4.5 g. yelloworange II [(RR' = (CH2)5] (VI), m. 130° (decomposition) (aqueous EtOH). VI (0.248 g.) in 30 cc. MeOH treated at room temperature with 20 cc. 0.1N AgNO3 gave 0.165 g. N, N-pentamethylene-N'-benzoylurea, m. 172° (decomposition) (dioxane-ligroine). I and 10 cc. Et20 treated with 2.6 g. cyclohexylamine in 20 cc. Et2O gave 2.9 g. II (R = cyclohexyl, R' = H) (VII), m. 150° (1:2 C6H6petroleum ether). I with 2.45 g. PhNH2 in 10 cc. dry Et20 stirred 10 min. at room temperature gave 3.0 g. sulfur yellow II (R = Ph, R'= H) (VIII), m. 214° (decomposition) (EtOH). VIII refluxed 0.5 hr. with 0.1N AgNO3-MeOH yielded 88% PhNHCONHBz. 2,3,6-Triphenyl-2H-1,3,5- thiadiazin-4-one (3.44 g.) in 50 cc. dioxane and 1 cc. H2O refluxed 5 min. gave 2.42 g. yellow VIII, m. 216° (decomposition). I (from 3.82 g. V) treated at 0° with 10 cc. dry AcOEt and then slowly with 3.38 g. Ph2NH in 10 cc. dry Me2CO and stirred 0.5 hr. at 0° yielded 30% PhCSNHCONPh2 (IX), m. 137° (decomposition) (petroleum ether). (0.332 g.) and 0.138 g. o-O2NC6H4NH2 in 7 cc. dry C6H6 heated 5 min. at 40° and kept at room temperature overnight yielded 0.19 g. II (R = o-O2NC6H4, R' = H) (X). I with 3.23 g. p-MeOC6H4NH2 in 30 cc. dry Me2CO yielded 4.84 g. bright yellow II (R = p-MeOC6H4, R' = H) (XI), m. 179° (decomposition). decomposed at about 200° with gas evolution and formation of a colorless solid, m. 230°. XI (1 g.), 0.007 mole Et3N, and 25 cc. dry AcOEt treated with

stirring at about 10° with 0.56 g. Br in 25 cc. dry AcOEt gave 0.5 g. light yellow XII (R = p-MeOC6H4), m. 155° (AcOEt). I with 3.62 g. o-O2NC6H4NH2 in 15 cc. dry Me2CO yielded 3.15 g. light brown-yellow X, m. 215° (decomposition) (C6H6). I and 4.6 g. 2,4-(O2N)2C6H3NH2 refluxed 1 hr. in 30 cc. dry Me2CO and stirred 20 min. yielded 0.9 g. II [R = 2,4-(O2N)2C6H3, R' = H], m. 225° (decomposition) (200:25 dioxane-H2O). I from 0.95 g. V treated dropwise with 0.59 g. p-H2NC6H4CN in 10 cc. absolute Me2CO and stirred 10 min. yielded 0.68 g. deep yellow II (R = p-NCC6H4, R' = H), m. 252° (decomposition) (PHCl). I from 1.91 g. V with 1.52 g. o-H2NC6H4CSNH2 in 10 cc. dry Me2CO gave 2.25 g. II (R = o-H2NCSC6H4, R' = H) (XIII), m. 198° (decomposition with formation of light yellow and red crystals). I from 1.9 g. V stirred 15 min. with 0.54 g. p-C6H4(NH2)2 in 10 cc. dry tetrahydrofuran yielded 1.05 g. yellow p-PhCSNHCONHC6H4NHCONHCSPh, decompose above 223° with the evolution of gas but without melting. I and 2.47 g. 2-aminopyridine in 15 cc. dry Me2CO stirred 15 min. gave 3.1 g. yellow II (R = 2-pridyl, R' = H), m 199° (decomposition) (AcOEt), which refluxed 4 hrs. with aqueous dioxane. qave a Sfree solid, m. 211° (decomposition). I with 2.5 g. 2-aminopyrimidine in 30 cc. dry Me2CO gave similarly 4.5 g. pink II (R = 2-pyrimidinyl, R' = H), m. 238° (decomposition) (HCONMe2). I with 4.65 g. 5-amino-3-phenyl-1,2,4thiadiazole in 30 cc. dry Me2CO stirred 15 min. gave 5.2 g. yellow II (R = 3phenyl-1,2,4-thiadiazol-5-yl, R' = H), m. 252° (decomposition) (HCONMe2tetrahydrofuran), which repptd. from AcNMe2 with petroleum ether gave orange prisms which change above 80° to the yellow form. I with 3.2 g. BzNH2 and 20 cc. dry Me2CO gave 1.3 g. IV, pink needles from C6H6, violet needles from Me2CO, m. 220 $^{\circ}$ (decomposition). PhCSNH2 (46 g.) in 400 cc. dry C6H6 refluxed 3 hrs. with 49 g. BzNCO yielded 80 g. IV. 2,6-Diphenyl-1,3,5-thiadiazin-4one (0.266 g.) in 5 cc. Me2CO heated briefly to 40° with a few drops H2O and 1 drop 2N HCl and kept 0.5 hr. at room temperature gave 0.27 g. IV. I and 3.6 g. BzNHNH2 in 25 cc. Me2CO yielded 2.6 g. yellow II (R = BzNH, R' = H) (XIV), m. 226° (decomposition) (C6H6). I from 2.5 g. V stirred 0.5 hr. with 1.57 g. PhCH:NNH2 in 10 cc. dry Me2CO yielded 0.82 g. light yellow II (R = PhCH:N, R' = H), m. 175° (decomposition). V (5 g.) and 4.0 g. H2NCH2CO2Et.HCl refluxed in methylcyclohexane gave 2.5 g. yellow PhCSNHCONHCH2CO2Et (XV), m. 138° (decomposition) (MeOH). XV (1 g.) and 10 cc. 4N NaOH heated about 10 min. at 40° and neutralized gave 0.85 g. light yellow PhCSNHCONHCH2CO2H, m. 258° with foaming (aqueous MeOH); it crystallized from aqueous MeOH with 0.5 mole H2O. I from 2.5 g. V with 0.66 g. N2H4.H2O in 15 cc. dry tetrahydrofuran yielded 1.2 g. yellowish XVI (R = R' = H) (XVII), m. 321° (aqueous EtOH). XIV (0.3 g.) and 1 drop Me2CO in 5 cc. 4N NaOH refluxed 10 min. and neutralized gave 0.15 g. XVII, m. 320-4°. I with 2.9 g. PhNHNH2 in 5 cc. dry Et20 at -20° gave 2.23 g. yellow precipitate which heated in AcOH gave with the elimination of H2S a mixture of XVI (R = Ph, R' = H) (XVIII) and XVI (R = H, R' = Ph) (XXIX) which fractionally recrystd. from aqueous AcOH gave 1.66 g. XIX, m. 235°, and 0.1-0.2 g. XVIII, m. 278° (partial decomposition). I from 1.91 g. V in 20 cc. methylcyclohexane refluxed 15 min. with 1.84 g. (PhNH)2 in 10 cc. absolute tetrahydrofuran gave 0.86 g. XVI (R = R' = Ph), m. 242° (decomposition) (EtOH). I with 3.2 g. PhC(:NH)NH2 in 20 cc. dry Me2CO refluxed 5 min. yielded 2.1 g. PhC(:NH)N:CPhNHCONHC(:NH)Ph (XX), m. 240-4° (decomposition) (AcNMe2-AcOEt). XX (about 0.5 g.) fused gave with the evolution of PhCN and NH3 2,6diphenyl-3,4-dihydro-1,3,5-triazin-4-one, m. 289° (C6H6N). I in 25 cc. methylcyclohexane with 5 g. PhC(:NH).NHPh in 20 cc. dry dioxane gave 2.4 g. 1,2,6-triphenyl-1,4-dihydro-1,3,5-triazin-4-one, m. 284° (decomposition) (tetrahydrofuran) with the formation of a solid, m. 232° with sublimation. XIII (0.78 g.) in 4 cc. dry Me2CO and 0.32 g. (COCl)2 in 10 cc. dry Me2CO gave at about 70° 0.63 g. red XXI, m. 163° (decomposition). IV (56.8 g.) in 100 cc. Me2CO and 2 l. 2N NaOH shaken 14 hrs. at room temperature and neutralized with AcOH yielded 30-1 g. lemon yellow III, m. 190° (decomposition) (AcOEtligroine). III (1.8 g.) in 10 cc. 2N NaOH treated gradually with 1.3 cc. 30% H2O2 gave XII (R = H), m. 204° (MeOH); it gives a blood red color with FeCl3-MeOH).

RN 5378-02-9 CAPLUS

CN Urea, 1,1'-p-phenylenebis(3-(thiobenzoyl)- (7CI, 8CI) (CA INDEX NAME)

L5 ANSWER 32 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1966:36325 CAPLUS Full-text

DN 64:36325

OREF 64:6778b-d

TI Acylisothiocyanates. VI. Reactions of bis(acyl isothiocyanates) with diamines

AU Li, Yung-Hsien; Chen, Yao-Tsu

CS Ind. Coll., Kansu, Peop. Rep. China

SO Gaofenzi Tongxun (1964), 6(3), 206-12 CODEN: KFTTAR; ISSN: 0453-2880

DT Journal

LA Chinese

acf. Sci. Sinica (Peking) 12, 143(1963); CA 52, 19993b. Bis(acyl isothiocyanates) reacted readily with diamines to form linear polymers of acylthioureas with the structure [R'NHCSNHCORCONHCSNH]n. Ten such poly(acylthioureas) were synthesized by the reactions of adipic, azelaic, and terephthalic diisothiocyanates with hydrazine, ethylenediamine, H2N(CH2)6NH, p-phenylenediamine, and benzidine. The structure of the polymers obtained was confirmed by elementary analysis, degradation examination, and uv and ir spectroscopy. These polymers were colored (yellow to orange) powders, sparingly soluble in common organic solvents, but readily soluble in HCONMe2 and cold concentrated H2SO4. The x-ray diffraction patterns showed that these polymers possessed fair crystallinity. The softening points of the polymers decreased with increasing length of the aliphatic C chain and increased when benzene nuclei were introduced into the chain. Four of these polymers had softening points >300°.

RN 70110-39-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CI) (CA INDEX NAME)

L5 ANSWER 33 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

DN 60:68587 OREF 60:12118e-g

TI Poly(acylthioureas)

AU Chen, Yao-Tsu; Li, Yung-Hsien

CS Univ. Lanchow, Peop. Rep. China

SO Kexue Tongbao (Chinese Edition) (1963), (10), 50-2 CODEN: KHTPAT; ISSN: 0023-074X

DT Journal

LA Unavailable

Diisothiocyanates of formula R(CONCS)2 (from diacyl chlorides and 2 moles AΒ NH4CNS) can add 2 moles of a primary amine, R'NH2, to form bis(acylthioureas), (R'NHCSNHCO)2R. For R' = Ph and R given, the m.ps. are: (CH2)4, $192-3^\circ$; p-C6H4 (I), 290°. If RCONCS (from RCOCl and 1 mole NH4CNS) was treated with diamines, R'(NH2)2, bis(acylthio-ureas) of type (RCONHCSNH)2R' were formed; e.g. for R = Ph and R' given, the m.ps. are: (CH2)6, 177-8°; p-C6H4, 237-8°. By hydrolysis with 10% NaOH, 80-90% of the original carboxylic acid and thiourea were recovered and identified by mixed-m.p. determination By keeping bis(acyl isothiocyanates) (3 kinds) and diamines (5 kinds) for 12 hrs. in anhydrous Me2CO, 10 poly(acyl-thioureas) were obtained containing the fundamental unit R'NH-CSNHCORCONHCSNH (R, R', m.p., and reduced viscosity at 30 \pm 1° in 0.5 g./ml. concentrated H2SO4 given): (CH2)4, (CH2)2, 185° (decompose), 0.10; (CH2)4, (CH2)6, 180° (decompose), 0.18 (infrared absorption bands at 5.58-6.1, 6.3-6.65, 7.8-8.0, 8.6, and 13.58 μ); (CH2)7, (CH2), 125- 9° , 0.10; (CH2)4, p-C6H4, m. >300°, 0, 20 (infrared absorption bands at 2-15 μ; ultra-violet absorption similar to that of I); (CH2)7, p-C6H4, 150-3°, 0.16; p-C6H4, -, m. >300°, 0.069; p-C6H4, (CH2)2, 210° (decompose), 0.12; p-C6H4, (CH2)6, 120-5°, 0.12; p-C6H4, p-C6H4, m.>300°, 0.11; and p-C6H4, p- ${\tt C6H4C6H4, m.>300°, 0.13.}$ The x-ray diagrams for most of the polymers indicate a crystalline state of linear order. The polymers are yellow or orange powders, insol. in most organic solvents, but readily soluble in HCONMe2 or concentrated H2SO4. Introduction of a benzene ring raises the softening point. The dielec. constant ranges from 1010 to 1011 ohm-cm.

Tollo-39-3, Urea, 1,1'-p-phenylenebis[3-benzoyl-2-thio-(preparation of)

RN 70110-39-3 CAPLUS

CN Benzamide, N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis- (9CÌ) (CA INDEX NAME)

L5 ANSWER 34 OF 34 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1961:111847 CAPLUS Full-text

DN 55:111847

OREF 55:21006d-f

TI Mono- and diisocyanates of p-cymene

AU Adellac, F.; Lora-Tamayo, M.; Soto, J. L.

CS Univ. Madrid

SO Anales real soc. espan. fis. y quim. (Madrid) (1960), 56B, 985-94

DT Journal

LA Unavailable

The reaction of phosgene with the appropriate amines was used to prep. the following isocyanates of cymene (substituents, b.p./mm., m.p., nD (t), and % yield given): 2-OCN, 76-7°/1, -, 1.5205 (22°), 70; 3-NCO, 76-7°/1, -, 1.5190 (22°), 60; 6-NO2, 2-NCO, 120-3°/1, 75°, 1.5425 (55°), 50; 2,6-(NCO)2 123-6°/2, 52-3°, 1.5517 (55°), 89; 2,5(NCO)2, 125-6°/2, 46-7°, 1.5394 (55°), 65; 3,5-(NCO)2, 110-12°/2, -, -, 81. The p-tolyl-, benzoyl-, phenylureas, and some of the methyl- and ethylurethans were prepared 2,3-Diamino-p-cymene (15 g.) in 300 ml. o-Cl2C6H4 treated with COCl2 several hrs., the mixture distilled, and cooled yielded 2-hydroxy-4-methyl-7-isopropylbenzimidazole, m. 260-1°, which with PCl5 yielded the 2-Cl derivative, m. 237-8°.

IT 124143-33-5, Urea, 1,1'-[2-isopropyl-5-methyl-p-phenylene]bis[3-benzoyl- 124143-34-6, Urea, 1,1'-(5-isopropyl-2-methyl-m-phenylene)bis[3-benzoyl- 124514-32-5, Urea, 1,1'-[2-isopropyl-5-methyl-m-phenylene]bis[3-benzoyl-

(preparation of)

RN 124143-33-5 CAPLUS

RN 124143-34-6 CAPLUS

RN 124514-32-5 CAPLUS

CN Urea, 1,1'-(2-isopropyl-5-methyl-m-phenylene)bis[3-benzoyl- (6CI) (CA INDEX NAME)

L6 5 L4 NOT L5

=> dis 1-5 bib abs

- L6 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 2005:980048 CAPLUS Full-text
- DN 143:359432
- TI Acyl Ureas as Human Liver Glycogen Phosphorylase Inhibitors for the Treatment of Type 2 Diabetes
- AU Klabunde, Thomas; Wendt, K. Ulrich; Kadereit, Dieter; Brachvogel, Volker; Burger, Hans-Joerg; Herling, Andreas W.; Oikonomakos, Nikos G.; Kosmopoulou, Magda N.; Schmoll, Dieter; Sarubbi, Edoardo; Von Roedern, Erich; Schoenafinger, Karl; Defossa, Elisabeth
- CS Scientific and Medical Affairs, Sanofi-Aventis Deutschland GmbH, Frankfurt am Main, D-65926, Germany
- SO Journal of Medicinal Chemistry (2005), 48(20), 6178-6193 CODEN: JMCMAR; ISSN: 0022-2623
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 143:359432
- AB Using a focused screening approach, acyl ureas have been discovered as a new class of inhibitors of human liver glycogen phosphorylase (hlGPa). The x-ray structure of screening hit 1 (IC50 = 2 μM) in a complex with rabbit muscle glycogen phosphorylase b reveals that 1 binds at the AMP site, the main allosteric effector site of the dimeric enzyme. A first cycle of chemical optimization supported by x-ray structural data yielded derivative 21, which inhibited hlGPa with an IC50 of 23±1 nM, but showed only moderate cellular activity in isolated rat hepatocytes (IC50 = $6.2 \mu M$). Further optimization was guided by (i) a 3D pharmacophore model that was derived from a training set of 24 compds. and revealed the key chemical features for the biol. activity and (ii) the 1.9 Å crystal structure of 21 in complex with hlGPa. second set of compds. was synthesized and led to 42 with improved cellular activity (hlGPa IC50 = 53±1 nM; hepatocyte IC50 = 380 nM). Administration of 42 to anesthetized Wistar rats caused a significant reduction of the glucagoninduced hyperglycemic peak. These findings are consistent with the inhibition of hepatic glycogenolysis and support the use of acyl ureas for the treatment of type 2 diabetes.
- RE.CNT 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L6 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 2005:495314 CAPLUS Full-text
- DN 144:254456
- TI Synthesis and characterization of novel multifunctional acylthiourea polymers
- AU Mao, Xue Pu; Huang, Jin Feng; Duan, Zhi Fang; Du, Zhi Yun; Huang, Zhi Shu; Ma, Lin; Gu, Lian Quan
- CS School of Chemistry and Chemical Engineering, Sun Yat-sen University, Guangzhou, 510275, Peop. Rep. China
- SO Chinese Chemical Letters (2005), 16(5), 609-612 CODEN: CCLEE7; ISSN: 1001-8417
- PB Chinese Chemical Society
- DT Journal
- LA English
- OS CASREACT 144:254456
- AB Several thiourea polymers have been synthesized through the reaction of diamine with 1,4- or 1,3-benzenedicarbonyl chloride and ammonium thiocyanate by solid-liquid phase transfer catalysis of polyethylene glycol-400 (PEG-400). The polymers were characterized and identified by elemental anal., IR, 1HNMR

and GPC. The multifunctional polymers have potential value as an ideal support for immobilized enzymes.

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 2004:432244 CAPLUS Full-text
- DN 142:155632
- TI Synthesis of novel bis-benzoylphenylurea chitin inhibitors
- AU Lin, Jun; Yang, Li-juan; Yan, Sheng-jiao; Li, Jun-feng; Liu, Fu-chu
- CS Department of Applied Chemistry, Yunnan University, Kunming, 650091, Peop. Rep. China
- SO Hecheng Huaxue (2004), 12(2), 117-119 CODEN: HEHUE2; ISSN: 1005-1511
- PB Hecheng Huaxue Bianjibu
- DT Journal
- LA English
- OS CASREACT 142:155632
- GΙ

- II
- AB Twelve novel bis-benzoylphenylurea chitin inhibitor derivs., I (R1 = C1, F; R2 = 4-C1, 2-C1, 4-Br) and II (R3 = H, CN, R4 = R5 = F, C1; R3 = H, CN, R4 = C1, R5 = H), have been synthesized in over 30 .apprx. 50% yield from chlorothalonil via sequential fluorine exchange, nitrile hydrolysis, decarboxylation and acylation reactions.
- RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L6 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 2004:60456 CAPLUS Full-text
- DN 140:128158
- TI Preparation of N-[(phenylamino)carbonyl]benzamides as glycogenphosphorylase-A inhibitors for the treatment of diabetes
- IN Defossa, Elisabeth; Kadereit, Dieter; Klabunde, Thomas; Burger,
 Hans-Joerg; Herling, Andreas; Wendt, Karl-Ulrich; Von Roedern, Erich;
 Schoenafinger, Karl
- PA Aventis Pharma Deutschland GmbH, Germany
- SO PCT Int. Appl., 75 pp.

CODEN: PIXXD2

DT Patent LA German FAN.CNT 1

r Auv.		PATENT NO.			KIND DATE			APPLICATION NO.					DATE					
PI	WO	WO 2004007437		A1	20040122		WO 2003-EP6934					20030630						
		W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
			PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,
			UA,	ŪG,	UΖ,	VC,	VN,	YU,	ZA,	ZM,	ZW							
		RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŬĠ,	ZM,	ZW,	AM,	AZ,	BY,
									ΑT,									
			FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
	CA	2493	373			AA		2004	0122	4	CA 2	003-	2493	373		2	0030	630
	ΑU	2003	2810	27		A1 20040202		AU 2003-281027					20030630					
	BR	2003	0125	93		Α		2005	0412	BR 2003-12593					20030630			
	ΕP	1523	471			A1		2005	0420		EP 2	003-	7403	86		20030630		
		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	ΗU,	SK	
	JP	2005	5324	02		Т2		2005	1027		JP 2	004-	5204	38		2	0030	630
		2004						2004	0506	1	US 2	003-	6169	59		2	0030	711
PRAI	DE	2002	-102	3137	1	Α		2002	0711									
	US	2002	-425	600P		P		2002	1112									
		2003				W		2003	0630									
os	MAI	RPAT	140:	1281	58													
GI																		

Title compds. I [W, X, Y = O, S; R9, R10, R11, R12 = H, halo, OH, etc.; R1, R2 = H, (un)substituted alkyl; R3, R4, R5, R6 = H, halo, OH, etc.; R7 = H, (un)substituted alkyl, e.g., OR13, NR14R15, etc.; R8 = NR18R19, OR20; R13 = H, alkyl, alkenyl, etc.; R14, R15 = H, (un)substituted alkyl; R18, R19 = H, alkyl, alkenyl, etc.; R20 = alkyl, alkenyl, alkynyl, etc.] and their pharmaceutically acceptable salts were prepared For example, condensation of benzamine II (Z = H), e.g., prepared from 2-chloro-4-fluorobenzamide in 2-steps, and carbonochloridic acid Me ester afforded benzamide II (Z = COMe) in 55% yield. In glycogenphosphorylase-A (GPa) inhibition assays, 23-examples of

compds. I, at 10 μ M, exhibited 48-100% inhibition of GPa activity, e.g., benzamide II (Z = COMe) displayed 53% enzyme inhibition. Compds. I were claimed useful as antidiabetic agents.

RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 2003:790618 CAPLUS Full-text
- DN 140:339042
- TI Synthesis and activities of aroyl(aryloxyacetyl) aryldithiourea derivatives as plant growth regulators
- AU Wu, Wei-lin; Ye, Wen-fa; Du, Zi-xiu; Wang, Yan-gang
- CS Huaihua Medical College, Huaihua, 418000, Peop. Rep. China
- SO Hecheng Huaxue (2003), 11(4), 310-314 CODEN: HEHUE2; ISSN: 1005-1511
- PB Hecheng Huaxue Bianjibu
- DT Journal
- LA Chinese
- OS CASREACT 140:339042
- AB By the use of solid-liq. phase transfer catalyst, 15 title compds. with diacylthiourea structure were synthesized from substituted aryloxyacetic acid or aromatic acid and aromatic diamine. For example, reaction of 3-MeC6H4CONCS, prepared from 3-methylbenzoic acid, with p-phenylenediamine gave 83% N,N'-[1,4-phenylenebis(iminocarbonothioyl)]bis[3-methylbenzamide]. The test of their biol. activities shows that most compds. have good plant growth regulating activities and a few of them are more active than indoleacetic acid.

SINCE FILE	TOTAL
ENTRY	SESSION
190.77	357.92
SINCE FILE	TOTAL
ENTRY	SESSION.
-29.25	-29.25
	ENTRY 190.77 SINCE FILE ENTRY

STN INTERNATIONAL LOGOFF AT 10:23:17 ON 11 OCT 2006